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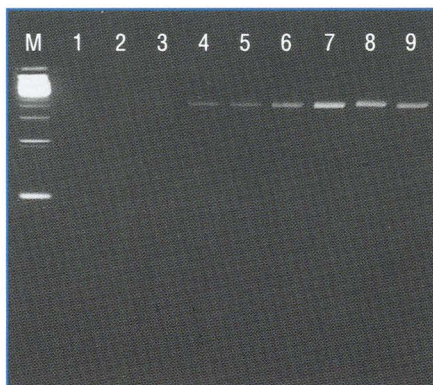
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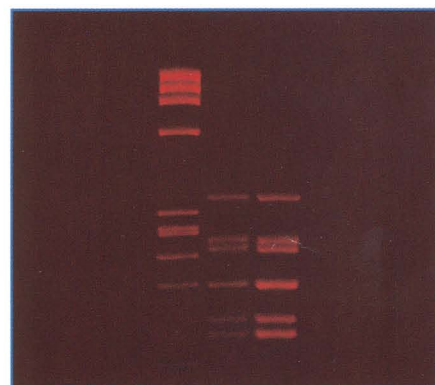
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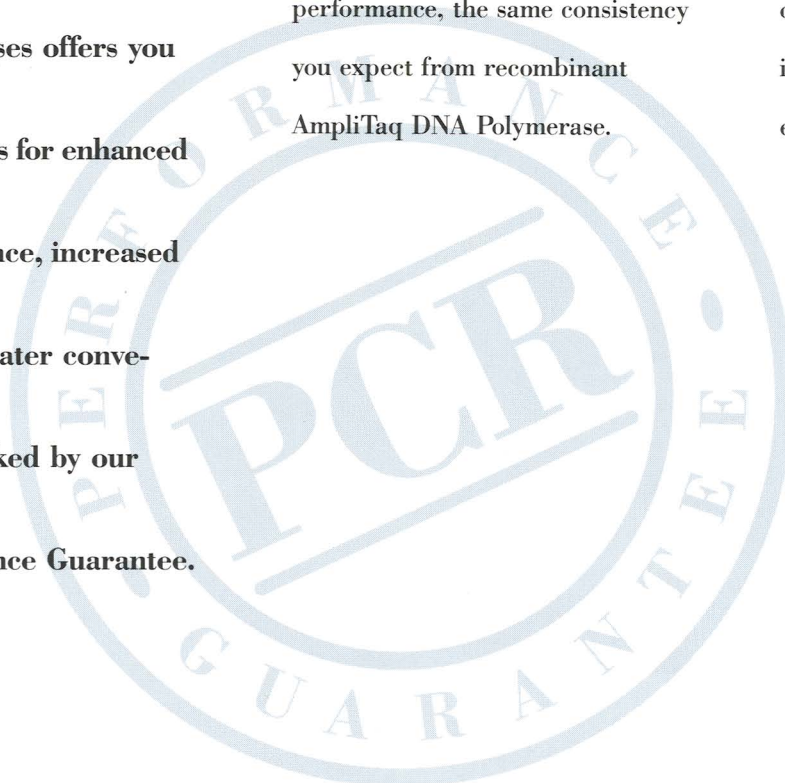
New AmpliTaq® DNA Polymerase, LD is ideal

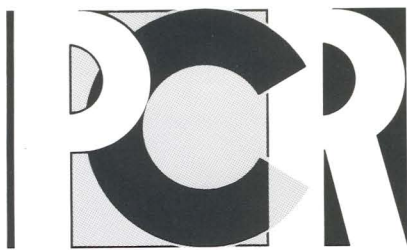
for low copy number amplifications of bacterial targets. A proprietary separation process has been used to reduce background DNA to fewer than ten copies. You'll find the same performance, the same consistency you expect from recombinant AmpliTaq DNA Polymerase.



AmpliTaq® DNA Polymerase, Stoffel Fragment meets special

PCR needs such as amplification of G+C rich templates and multiplex PCR. AmpliTaq DNA Polymerase, Stoffel Fragment features increased thermal stability, optimal activity over a broad range of magnesium ion concentrations and lack of 5'-3' exonuclease activity.





METHODS • AND • APPLICATIONS

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